Application No. 09/328,742

Amendment Dated: December 1, 2008 Reply to Office Action of: May 30, 2008

REMARKS

No claims are added. Claims 1-23 and 26 are canceled. Claims 24 and 28 are amended. Claims 24, 25 and 27-34 are currently pending.

Rejection of claims 23-25, 28-30, 33 and 34 as indefinite

The Examiner rejects claims 23-25, 28-30, 33 and 34 under 35 U.S.C. 112 as indefinite because the previous amendment did not indicate removal of the entire structure AA-CO-NH-CH(CH3)CH2OH in independent claim 28. Applicant believes that the Examiner actually meant that the enter structure "AA-CO-O-C(CH2OH)2" was not removed in the prior amendment. Regardless, Applicant has amended claim 28 to indicate removal of both structures, thus obviating the rejection.

Rejection of claims 28-30, 33 and 34 as anticipated

The Examiner also rejects claims 28-30, 33 and 34 under 35 U.S.C. 102(b) as anticipated by a reference to Abadji et al. (R-Methanandamide: A Chiral Novel Anandamide Possessing Higher Potency and Metabolic Stability, J. of Medicinal Chem., 1994, Vol. 37, no. 12, pp. 1889-1893).

Accordingly, Applicant has amended independent claim 28 by removing the compound AA-CO-NH-CH(CH3)CH2OH and respectfully requests that the Examiner withdraw his rejection.

Rejection of claims 25, 27, 28 and 31-34 as anticipated

The Examiner rejects claims 25, 27, 28 and 31-34 under 35 U.S.C. 102(b) as anticipated by a reference to Calignano et al. (Potentiation of anandamide hypotension by the transport inhibitor, AM404, European J. of Pharm., 1997, vol. 337, pp. R1-R2).

As argued in the Response to Office Action filed on August 15, 2006 and accepted by the Examiner in the Office Action of November 22, 2006, Calignano

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is NOT prior art to the present Application. Regardless, Applicant reiterates his remarks from the August 15, 2006 Response as follows:

U.S. Provisional Patent Application No. 60/088,568 supports the use of AM404

The present Application properly claims priority from U.S. Provisional Patent Application No. 60/088,568. The '568 application discloses the structure of compound AM404 and its use as an anandamide transport inhibitor. The '568 application also provides guidance on where to find synthesis information for compound AM404 (see page 9).

• Calignano is not prior art under 35 U.S.C. 102(b)

The present Application properly claims priority from the '568 provisional application filed June 9, 1998. Consequently, the proper effective filing date for the present Application is June 9, 1998 and the proper 102(b) bar date is June 9, 1997.

Calignano was published August 21, 1997. Consequently, Calignano is not prior art under 35 U.S.C. 102(b). At best, Calignano may be prior art under 35 U.S.C. 102(a).

• A 35 U.S.C. 102(a) reference may be overcome by submission of a declaration under 37 CFR 1.131

Rejections under 35 U.S.C. 102(a) can be overcome by submitting an affidavit or declaration under 37 CFR 1.131 showing prior invention (see MPEP § 706.02(b)). Enclosed herewith are copies of the Declaration of Alexandros Makriyannis Under 37 CFR 1.131 and Exhibits A-E, as originally submitted with the Response to Office Action of August 15, 2006.

As attested to in the Declaration, Applicant's invention of the recited subject matter predates the publication date of the Calignano reference.

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Consequently, Calignano is not prior art under 35 U.S.C. 102(a) and Applicant respectfully requests that the Examiner withdraw the rejection of claims 25, 27, 28 and 31-34 as anticipated by Calignano.

Rejection of claims 23 and 24 as obvious

The Examiner rejects claims 23 and 24 under 35 U.S.C. 103(a) as obvious in view of Abadji. Claim 23 is canceled, thus obviating its rejection. Additionally, claim 24 is amended to remove the stereoisomers of the compound disclosed by Abadji. Consequently, claim 24 is allowable.

For the reasons stated herein, the pending claims are not anticipated or obvious. Applicant respectfully requests that the Examiner withdraw his rejections and pass the pending claims to issue.

Respectfully submitted,

ALEXANDROS MAKRIYANNIS et al.

By:

Alexander E. Andrews Registration No. 62,205 Alix, Yale & Ristas, LLP Attorney for Applicant

Date: /2.01.2002 750 Main Street, Suite 1400 Hartford, CT 06103-2721 (860) 527-9211 Our Ref: UCONAP/141/US

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DECLARATION OF ALEXANDROS MAKRIYANNIS UNDER 37 C.F.R. 1.131

I, Alexandros Makriyannis, hereby declare:

- 1. I am a co-inventor named in U.S. Patent Application No. 09/328,742. I have reviewed this application.
- 2. I am a co-author of a printed publication article titled "Functional Role of High-Affinity Anandamide Transport, as Revealed by Selective Inhibition" published in SCIENCE, volume 277, pages 1094 -1097 and dated August 22, 1997. I have reviewed this article.
- 3. Attached hereto are documents containing facts showing the preparation of N-(4-hydroxyphenyl)arachidonylamide (compound AM404) in the United States before August 1, 1997. The dates on all documents have been redacted, which dates are prior to August 1, 1997.
- 4. Also attached hereto are documents containing facts showing that N-(4-hydroxyphenyl)arachidonylamide (compound AM404) was tested in the United States before the August 1, 1997. The dates on all documents have been redacted, which dates are prior to August 1, 1997.
- 5. Exhibit A is a photocopy of pages of a laboratory notebook illustrating an experiment testing compound AM404 for anandamide uptake in cells. These pages have been labeled with numbers 000001 to 000004 for convenience.
- 6. Exhibit B is a photocopy of pages of a laboratory notebook illustrating another experiment testing compound AM404 for anandamide uptake in cells. These pages have been labeled with numbers 000005 to 000008 for convenience.
- 7. Exhibit C is a photocopy of pages of a laboratory notebook illustrating another experiment testing compound AM404 for anandamide uptake in cells. These pages have been labeled with numbers 000009 to 000012 for convenience.
- 8. Exhibit D is a photocopy of pages of a laboratory notebook illustrating another experiment testing compound AM404 for anandamide uptake in cells. These pages have been labeled with numbers 000013 to 000016 for convenience.
- 9. Exhibit E is a photocopy of pages of a laboratory notebook illustrating another experiment testing compound AM404 for anandamide uptake in cells. These pages have been labeled with numbers 000017 to 000020 for convenience.
- 10. Exhibits A-E illustrate that administration of compound AM404 to cells inhibits transport of anandamide in those cells.

I hereby further declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

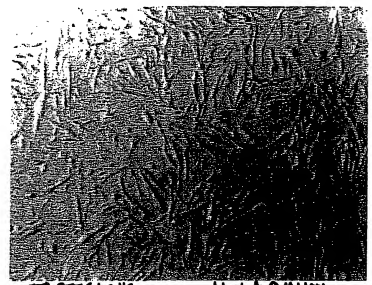
Date

Alexandros Makriyannis

EXHIBIT A

Used CCF. STTGI cells for an uptake experiment following the uptake protocol 1:00 & utilizing the multichanted Note: For this experiment we are lowering prettor.

The CD of unlabelled amandamod from 100mm to 30mm.



Compound tested: AMHOY
cells 5 daws in
culture sine replating
(5th replating)

Plate SIT assay

30 nm mabelled anordamide

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CCF-STTGI	295-297	11	+	18		0.1%		
	Z98.300	11	+	11	+	0.03	MM	DW HOH
	301 - 303	11	+	11	+	0.1	MM	404 MA
	304-306	11	}-	11	+	0.3	MΜ	POH MA
	367 - 309	11	+	11	+	1.0	MM	AM 404
	310-312	11	+	11	+	3.0	MM	FM HDH
	313-315	۱۸ ،	+	"	+	10.0	MM	HOP MA
	316.38	11	+	11	۲	30.0	MM	POH MA

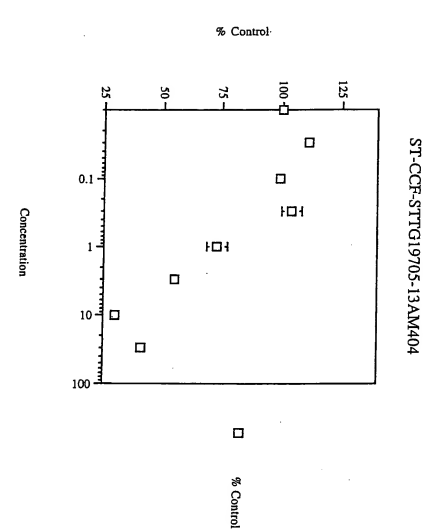
Preincubation (w) the same drug CI's as incubation) was carried out for 13 min. Incubation for 4 minutes. Cells were detached from the plates by sonicating in a water both for 30 seconds.

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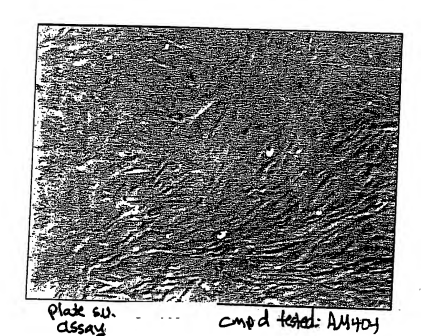
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EXHIBIT B

Used CCF-STTG1 cells for an uptake experiment following the uptake protocol 1.00



Completed: AMHOY

Cells in culture 7

days since replating.

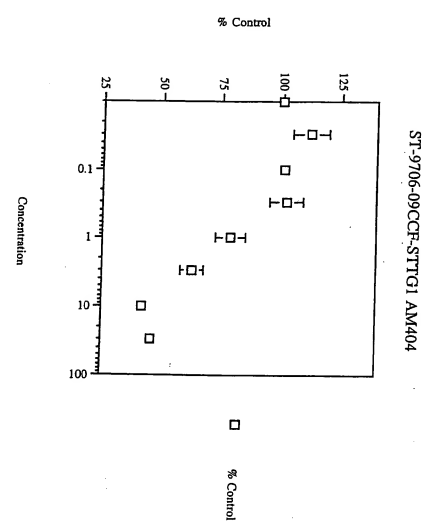
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S65 - 567) p	+	77	+	0.1 MM
<i>5</i> 68 - 570	11	+	**	+	D.3 MM
S71-573	, (1)	+	11	*	1.0 MM
574 - 576	11	+	11	+	3.0 MM
577-579	\ t	+	11	4	MM 0.01
580 - 58Z	γt	+	11	+	30.0 MM

Pre-incubation time: 13 min. incubation time: 4 min.

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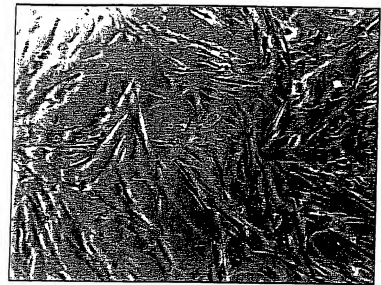
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EXHIBIT C

used (CF. STTGI cells for an uptake experiment following whe uptake protocol 1:00



CCF-STTE: AMYOU assay

Completed: AM 404 Cells in where 4 days Since replating.

CCF-STTG-1	30mm un	would o	mandemide		
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661-663	7)	+-	11		POPMA MU 1.0
664-6 66	11	+	11		O.S MM AM YOU
667-669	11	+	1 (1.0 MM BU 404
670-672	//	+	. 11		3.0 MM AM 404
673-675	11	4	17		10.0 MM AM 404
676-678	T.	+	()	+-	30.0 MM AM 404

Preinculation time: Bmin. Incubation time: 4 min.

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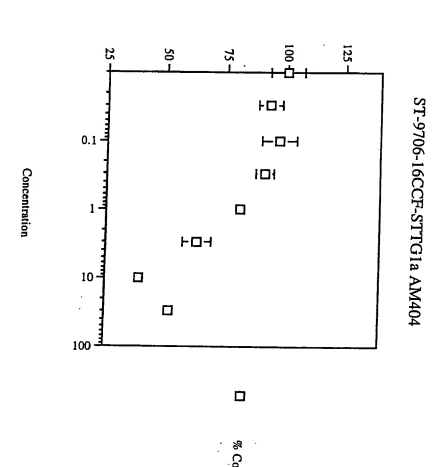
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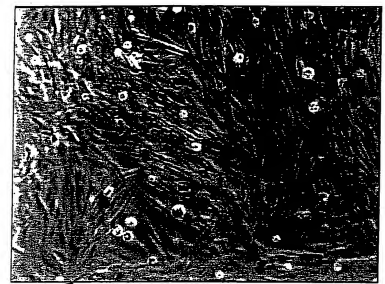
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1515.24 3.9363e+04 3.2403e+05 3.327e+04 3.8571e+05	2024.4/ 2422.05	2624 17	PIBO MPL	D	0.3	0
3 3 1398.37 1284.66 1.4114e+05.8921e+05 .6149e+05.8977e+05 .7349e+05.2094e+05 .1768e+05.6516e+05	2422.05	.01316+046	Haw Data	7.0 D		7
3 1284.66 8921e+06 9877e+06 2094e+06 6516e+06	2225.09	.4449e+045	Raw Data	3.0 In	-	
3 318.32 6184e+0.5. 8923e+0.6. 6950e+0.6. 8031e+0.5.	551.35	75536+045	Raw Data	10.0 In	G	
3 8.32 878.73 4e+0.5.2854e+0.4 3e+0.5.0417e+0.4 3e+0.5.350e+0.4 1e+0.5.8316e+0.4	1522 01	4	3	30.0 In	Į	
7962.80 5397.00 6894.00	755.33	1	Control	Control	-	
492.64 3 284.43 4112.44 6560.22 4837.00 5822.00	5336.33	Raw Data	0.03 c	٥		
826.92 3 477.43 4276.97 8385.69 5413.00 7017.00	6331.33	Raw Data	0.1 c	~		

EXHIBIT D

Utilized CCF.STTG1 cells for an uptake exporment following the uptake protocol 1.00



SCF-STTGI

assay:

Compa lested: AMYOY

(place reversed)

Compad tested: AM404

Cells in culture 5 days since replating.

For an experiment, we reversed the orientation of the plate, to assure that different positioning of the dilutions gave the usual result.

usual position:

Control	1.0
0.03	3.0
0.1	10.0
0.3	30.0

Position this I

1.0
3.0
10.0
30.0

F-STTG1 30nM unlabelled anardamide+

849-851	0.45mm	[3H] aran	damide	+	Q1% DMSD	\ <u></u>
852-854	11	+	1,	7	0.03 mM A	M404
855-857	11	+	11		0.1 MM A.	
828.860	11	+	11		0.3 jum A	
861-863	11	+	**		1.0 MM A)	
844-866	**	+	**		3.0 MM A	
867-867	11	+	W		10-0 MM A	
868-870	**	+	1.		300 MM A.	

Are-incubation time: 13 min. Incubation time: 4 min.

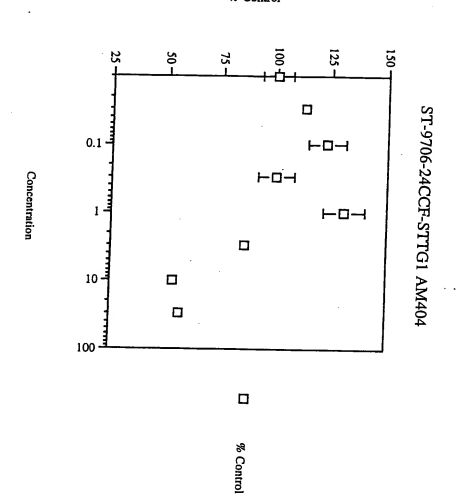
Cells detached by Sonicating for 60 seconds

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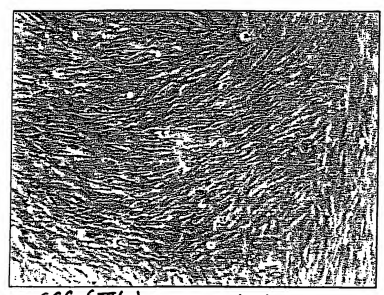
AM404

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4269.00	3875.00	4658.45	3586.89	124.51	3	215.66	4122.67	Haw Data	3.0 с	z
3594.00	2919.00	4082.11	2399.89	195.47	အ	338.57	3241.00	Raw Data	10.0 c	0
3822.00	3430.00	4099.73	3103.61	115.75	ယ	200.48	3601.67	Raw Data	30.0 с	ס

	Maximum	MINIMIM	IBX			Sample size	1	меап	or Mean	Column Tille	Column ID	I III e
	5.31200+04	5.0669e+04.5905e+04.9536e+04.3426e+04.82020+04.1509e+04.3875e+04	5.5584e+04.1976e+04.8844e+04.0555e+04.3809e+04.3809e+04.1500000000000000000000000000000000000	9.8865e+04.3663e+04.6072e+04.0802e+04.6116e+04 7955e+04 70	780.79	3	1352.36	5.2225e+04.7820e+04.2458e+04.5679e+04.8962e+04.0772e+04.02	Raw Data	Control in	A	CCt-S11G1 Uptake & Inhibition with
	.90010+04	.5905e+043	.19760+04	.36630+043	966.03	3	1673.21	.78206+04	Raw Data	0.03 in	В	ake & Inhibitio
	.43710+04	.9536e+043	.88440+04	.60720+043	1484.12	အ	2570.58	.24580+043	Raw Data	0.1 in	C	n with
	7022404	3426e+0A	.0555e+04	.08020+043	1133.28	ယ	1962.90	.5679e+04	Raw Data	0.3 in	D	
11136+043		82020100	38096+04	6116e+00	893.91	. د	1548.31	99626+04	Raw Data	1.0 ln		
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01730+04.	83636+041.	10049+041	70118+0/3	06.226	500	900.11	0000+04	Daw Dala	Die or	5		
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9213.00	7367 00	05270+04	5710.79	559.66 O	<u>သ</u>	969.36	8119.00	Raw-Data	e.i.	~		

EXHIBIT E

Used CCF-STTGI cells for an uptake experiment following the uptake protocol 1:00:



S.v. assay

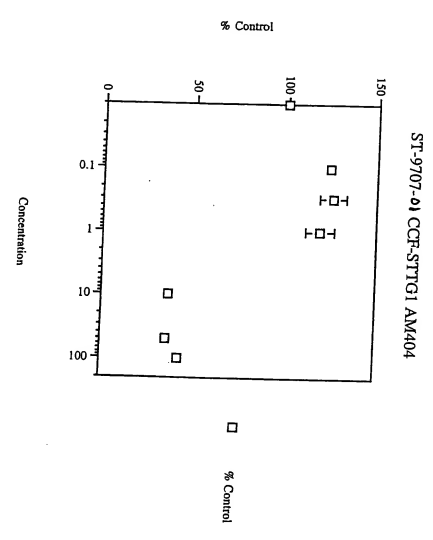
Completed: AM404 Cells in culture 6 days since replating.

CCF-STTG		30nM U	nlabelled_au	woda	mide	_	
	991-993	+ 0.45	NM [3H] a	nand	amide t	0.1% D	450
	994 - 996	15	+	A	+	0.1 mM	HOPMA
	997-999	11 ,	7-	11	+	0.3 MM	POPMA
•	1000 - 1002	11	+	**	+	MM OIL	404MA
	D03 - 1005	11	+	11	+	3.0 LM	AM 404
	1006-1008	11	+	11	+	Mu 0.01	
	1101-2001	11	+-	7)	+	50.0 mM	
	1012-1014	11	+	n	+	100.0 MM	AMHOY

Pre-incubotion Home. 13 mm. Incubation time: 4 mmin Cells detached by somicating 60 sciences

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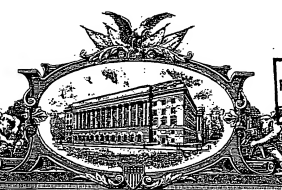
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CCF-STTG1 AM404

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Column ID Column Title Raw or Mean SD Sample size SEM 95% CI min 95% CI max Minimum Maximum
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CCI-STTG1 Uptake & Inhibition with A B Control in O.1 in 0.3 Rew Data Raw
CCI-STTG1 Uptake & inhibition with A B C C Control in O.1 in O.3 in I.0 in Raw Data
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CCI-STTG1 Uptake & Inhibition with CCI-STTG1 Uptake & Inhibition with E F G H I A B C D E F G H I Control in 0.1 in 0.3 in 1.0 in 3.0 in 10.0 in 50.0 in 100.0 in Control c B Control in 0.1 in 1.0 in 3.0 in 10.0 in 50.0 in 100.0 in Control c B Control in Raw Data
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TO ALL TO WHOM THESE: PRESENTS SHALL COME:
UNITED STATES DEPARTMENT OF COMMERCE
United States Patent and Trademark Office

August 30, 1999

THIS IS TO CERTIFY THAT ANNEXED HERETO IS A TRUE COPY FROM THE RECORDS OF THE UNITED STATES PATENT AND TRADEMARK OFFICE OF THOSE PAPERS OF THE BELOW IDENTIFIED PATENT APPLICATION THAT MET THE REQUIREMENTS TO BE GRANTED A FILING DATE UNDER 35 USC 111.

APPLICATION NUMBER: 60/088,568

FILING DATE: June 09, 1998

PCT APPLICATION NUMBER: PCT/US99/12900



By Authority of the COMMISSIONER OF PATENTS AND TRADEMARKS

R. BLAKENEY
Certifying Officer

PRIORITY DOCUMENT

SUBMITTED OR TRANSMITTED IN COMPLIANCE WITH RULE 17 1(a) OR (b)



Aprov

PROVISIONAL APPLICATION FOR PATENT COVER SHEET

This is a request for filing a PROVISIONAL APPLICATION under 37 CFR 1.53 (b)(2).

		Docket Number	UCON/1	40/US	Type a plus sq	pn (+1	+					
	INV	ENTOR(s)/APPLIC	ANTIel									
LAST NAME	FIRST NAME	MIDDLE INITIAL		ICITY AND EITHE	R STATE OR FOI	REIGN CO	UNTRY					
Makrıyannis	Alexandros		Watert	own, MA,	USA							
Lin	Sonyuan		Storrs,	CT, USA								
TITLE OF THE INVENTION (280 characters max)												
Anandamide Transporter Inhibitor Medications												
CORRESPONDENCE ADDRESS.												
James E. Alix, Esq. Alix, Yale & Ristes, LLP 750 Main Street Hartford STATE CT ZIP CODE 06103-2721 COUNTRY U.S.A.												
STATE CT	ZIP CODE 06103-27	721 (OUNTRY	U S.A.								
ENCLOSED APPLICATION PARTS (check all that apply)												
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Respectfully submitted, SIGNATURE S. CO.: Date JUNE 9, 1998 TYPED or PRINTED NAME James E. Alix REGISTRATION NO. 20,736 Applitional inventors are being named on separately numbured sheets attached hereto.												
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INVENTION DISCLOSURE

Anandamide Transporter Inhibitor Medications

Disclosure No.

Page 1

COMPLETE DESCRIPTION OF THE INVENTION: Use additional pages, if necessary, and attach any relevant sketches, diagrams, drawings, photographs or other illustrative material. ALL ATTACHED MATERIALS MUST BE SIGNED AND DATED BY EACH INVENTOR AND WITNESSED. Description may be by reference to a separate document sucl as a publication, manuscript, preprint or report. Such documents must be attached.				
A carrier protein that transports extracellular anandamide across the cell membrane has been shown to be present in rat neurons and astrocytes. This carrier protein or anandamide transporter is believed to be responsible for the inactivation of anandamide, an endogenous cannabinoid for central cannabinoid receptors. Thus, anandamide released from neurons on depolarization is rapidly transported back into the cells and subsequently hydrolyzed by an amidase thereby terminating its biological actions. Anandamide transporter is a potential therapeutic target for the development of useful medications.				

We have discovered a phenolic analog of anandamide namely N-(4-hydroxyphenyl)arachidonylamide (AM404) which inhibits the transport of anandamide across the cell membranes. AM404 does not activate cannabinoid receptors or inhibit anandamide hydrolysis per se. However, it does potentiate receptor-mediated anandamide responses by preventing anandamide reuptake.Continued on Supplement Page

(2) NOVEL FEATURES: Clearly specify the novel aspects of your invention. Compared to present technology, how is your invention different?

AM404 is a potent inhibitor of anandamide transport and it is the only compound known todate that competitively inhibits anandamide reuptake.

What deficiency in the present technology does your invention improve upon? Is it more effective? cheaper? superior in other ways?

Present cannabinoid drugs are targeted towards cannabinoid receptors (CB1 and CB2) and anandamide amidase enzyme. AM404 described in this invention targets a novel protein called anandamide transporter.

(3) STAGE OF DEVELOPMENT: Cite your specific results to date demonstrating that your concept is valid. Has your work included laboratory studies? Pilot-scale experiments? Construction and testing of a prototype?

AM404 inhibited accumulation of anandamide in rat neurons and astrocytes with an IC₅₀ of 1µM for neurons and 5µM for astrocytes. In addition, AM404 potentiated and prolonged receptor-mediated effects of anandamide such as vasodialation. These experiments further support that AM404 is an inhibitor of anandamide transport.

Inventor(s) 1! Helli yany Date 5/12/98 Disclosed to and Understood by:

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SUPPLEMENT PAGE

INVENTION DISCLOSURE FORM

Disclosure I	٧n		

Continued from page 1, item 1:

Structural formulas for AM404 and anandamide are shown below.

AM404 and its analogs are potential drug candidates for the treatment of ailments related to the cannabinoid system. Potential therapeutic uses of AM404 are pain alleviation (analgesia), treatment of cardiovascular diseases and blood pressure disorders.

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INVENTION DISCLOSURE

Page 2

Disclosure No._____

(4) VARIATIONS OF THE INVENTION: Discuss all alternate forms that you can foresee for this invention, whether or not you have evaluated them to date. (For example, chemical inventions should consider analogs and derivatives.)

AM404 was first synthesized in March 1993 and tested in July 1997 as anandamide transport inhibitor

(6) INVENTOR'S PUBLICATION PLANS: Please list all your publications — theses, reports, pre-prints, abstracts, papers, etc. that pertain to the invention. Include publication dates. Also, include manuscripts for publication (submitted or not), news releases, and internal publications. Enclose copies of all the above items with this disclosure.

Beltramo, M.; Stella, N.; Calignano, A.; Lin. S.; Makriyannis, A.; Piomelli, D. Functional Role of High Affinity

Anandmide Transport Inhibitor, as Revealed by Selective Inhibition. Science 1997, 277, 1094. (included) - CD&

BioWorld Today, Volume 8(162), August 21, 1997.

(7) PRIOR DISCLOSURE: Please give the details (date, place and circumstances) of any oral or written disclosures of all or part of this invention. If disclosed to specific individuals, give their names. Include professional meetings and conferences. Has this invention or a product resulting from this invention been offered for sale or license? Have any samples related to this invention been distributed?

No prior disclosure

Inventor(s) 1

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INVENTION DISCLOSURE

Page 3

Disclosure No.

SUPPORTING INFORMATION

(1) PRIOR KNOWLEDGE AND COMPETING RESEARCH AND DEVELOPMENT: Please list all publications and patents by the inventor or others that relate to the invention. The inventor should thoroughly search the published literature and review closely related patents.

Publications by the researchers:

1) Calignano, A.; La Rana, G.; Beltramo, M.; Makriyannis, A.; Piomelli, D. Potentiation of Anandamide Hypotension by the Transport Inhibitor, AM404. Eur. J. Pharmacol. 1997, 337, R1-R2. 2) Calignano, A.; La Rana, G.; Makriyannis, A.; Lin, S.; Beltramo, M.; Piomelli, D. Inhibition of Intestinal Motility by Anandamide, an Endogenous Cannabinoid. Eur. J. Pharmacol. 1997, 340, R7-R8.

List any known research groups currently engaged in research and development in this area. Include both academic and industrial researchers.

None

(2)ALTERNATE TECHNOLOGY: Describe any known alternate technologies that accomplish the same or similar purposes as this invention. List companies and products that currently use these alternate technologies.

None

(3) COMMERCIAL APPLICATION OF THE INVENTION: List all products, processes, devices, equipment, etc., to which your invention could be applied or which could result directly from your invention. Can these applications be developed in the near term (within two years) or the long term (more than two years)?

Medication to alleviate pain and treatment of cardiovascular diseases.

Long term development

What firms or types of firms do you think may be interested in the invention? Why? Name companies and specific persons if possible. Especially list companies with which you have had direct contact.

Pharmaceutical and biotech companies

(4) RESEARCH AND DEVELOPMENT PLANS: What additional research is needed to complete development and testing of the invention? Are you actively pursuing the needed work? Under whose sponsorship? About how long will this work take? What additional research support, if any, is needed for these efforts?

6

IDENTIFICATION AND FUNCTIONAL ROLE OF HIGH AFFINITY ANANDAMIDE TRANSPORT

M. Beltramo*, N. Stella*, A. Calignano*, S.Y. Lin*, A. Makriy unis* and D. Piomelli*. *The Neurosci. Inst. San Diego, CA 92121, #Sch. of Pharm. Univ. of Naples, Italy 80131, and *Sch. of Pharm. Univ. of Connecticut, Storrs, CT 06269.

Anandamide (arachidonylethanolamide) is an endogenous lipid that activates brain cannabinoid receptors. Two main pathways have been proposed for anandamice inactivation: cellular uptake and enzymatic degradation. In the present study we identified and the acterized pharmacologically a high affinity anandamide uptake system in neurons and astrocytes. Exogenous [3H]anandamide (spec. rad.: 221 Ci/mmol) is rapidly cleared (11/2=4 minutes) from the media of neurons or astrocytes in cell culture through a saturable, temperature-dependent and sodium-indepent ent transport system. This uptake displays high affinity for [3H]anandamide (neurons: Km 1.2 microM; astrocytes: Km 0.32 microM). Competition experiments with fatty acid derivatives, arachidonic ac d, or palmitoylethanolamide proved its specificity. Screening of lipid uptake blockers and anandamide : nalogs led to the identification of a compound N-(4-hydroxyphenyl) arachidonylamide (AM404) which is potent and specific in inhibiting anandamide transport, but does not activate CB) cannabinois receptors and does not inhibit anandamide degradation. In cultures of cortical neurons, concentrations of anandamide higher than 0.3 microM are necessary to activate CB1 cannabinoid receptors and to revert forskolin-induced adenylyl cyclase activity. In the presence of AM404 (10 microM) the potency of anandamide is greatly increased. By contrast, AM404 has no effect on adenytyl cyclase activity when applied alone (10 microM), and does not potentiate adenylyl cyclase activity inhibition elicited by the CB1 receptor agonist WIN-55212-2 (100 nM) or by glutamate (3 microM). The hot-plate model of analgesia in the mouse was used to test the functional role of anandamide transport in vivo. Intravenous (i.v) administration of anandamide (20 mg/kg) induces a modest, but significant, analgesia which disappears 60 minutes after the injection and is prevented by SR-141716 (1 m 1/kg, i.p.). Administration of AM404 (10 mg/kg, i.v.) has no antinociceptive effect per se within 60 t unutes of injection, but significantly enhances and prolongs anandamide-induced analgesia. The identification in neural cells of a highaffinity [3H]anandamide transport system and the discovery of selective transport blockers should be important to understand the physiological role of the endogenous cannabinoid system. In light of the multiple behavioral effects of cannabinoid receptor activation, these inhibitors might also open novel therapeutic avenues for the treatment of psychiatric and neurologi :al disorders.

Work at The Neurosciences Institute was supported by Neurosciences Resear: h Foundation which receives major support from Sandoz Pharm. S.Y. Lin and A. Makriyannis were supported by a grant (DA-3801) from NIDA.

Functional Role of High-Affinity Anandamide Transport, as Revealed by Selective Inhibition

M. Beltramo, N. Stella, A. Calignano, S. Y. Lin, A. Makriyannis, D. Piomelli*

Anandamide, an endogenous ligand for central cannabinoid receptors, is released from neurons on depolarization and rapidly inactivated. Anandamide inactivation is not completely understood, but it may occur by transport into cells or by enzymatic hydrolysis. The compound N-(4-hydroxyphenyl)arachidonylamide (AM404) was shown to inhibit high-affinity anandamide accumulation in rat neurons and astrocytes in vitro, an indication that this accumulation resulted from carner-mediated transport. Although AM404 did not activate cannabinoid receptors or inhibit anandamide hydrolysis, it enhanced receptor-mediated anandamide responses in vitro and in vivo. The data indicate that carrier-mediated transport may be essential for termination of the biological effects of anandamide, and may represent a potential drug target.

Anandamide (arachidonylethanolamide) is an endogenous lipid that activates brain cannabinoid receptors and mimics the pharmacological effects of A9-tetrahydrocannabinol, the active principle of hashish and marijuana (1). In humans, such effects include euphoria, calmness, dream states, and drowsiness (2). Depolarized neurons release anandamide (3) through a mechanism that may require the calcium-dependent cleavage of a phospholipid precursor in neuronal membranes (4). Like other modulatory substances, extracellular anandamide is thought to be rapidly inactivated, but the exact nature of this inactivating process is still unclear. A possible pathway is hydrolysis to arachidonic acid and ethanolamine. catalyzed by a membrane-bound facty acid amide hydrolase (FAAH) highly expressed in rat brain and liver (5). Nevertheless, the low FAAH activity found in brain plasma membranes indicates that this enzyme may be intracellular (5), a possibility that is further supported by sequence analysis of rat FAAH (6). Although anandamide could gain access to FAAH by passive diffusion. the transfer rate is expected to be low because of the molecular size of this lipid mediator (7). In that other lipids including polyursaturated fatty acids and prostaglandin E2 (PGE2) enter cells by carrier-mediated transport (8, 9), it is possible that anandamide uses a similar mechanism. Indeed, the existence of a rapid, saturable process of anandamide accumulation into neural cells has been reported (3). This

accumulation may result from the activity of a transmembrane carrier, which may thus participate in termination of the biological actions of anandamide. Accordingly, we developed drug inhibitors of anandamide transport and investigated their pharmacological properties in cultures of rat cortical neurons or astrocytes.

The accumulation of exogenous (3H)anandamide by neurons or astrocytes fulfills several criteria of a carrier-mediated transport (Fig. 1) (10). It is a rapid process that reaches 50% of its maximum within about 4 min (Fig. 1A). Furthermore, (3H)anandamide accumulation is temperature-dependent (Fig. 1A) and saturable (Fig. 1, B and C). Kinetic analyses revealed that accumulation in neurons can be represented by two components of differing affinities (lower affinity: Michaelis constant, $K_m = 1.2 \mu M$, maximum accumulation rate. max, = 90.9 pmol/min per milligram of protein; higher affinity: $K_m = 0.032 \mu M_s V_m$ 59 pmol/min per milligram of procein) (Fig. 1B). The higher affinity component may reflect a binding site, however, as it is displaced by the cannabinoid receptor antagonist, SR-141716-A (100 nM) (11). In astrocytes, (3H)anandamide accumulation is represented by a single high-affinity component (K = 0.32 µM, V_{max} = 171 pmoVmin per milligram of protein) (Fig. 1C). Such apparent K_m values are similar to those of known neurogransmitter uptake systems (12) and are suggestive therefore of high-affinity carrier-mediated

To characterize further this putative anandamide transport, we used cortical ascrocytes in culture. As expected from a selective process, the temperature-sensitive component of (PH)anandamide accumulation was prevented by nonradioactive anandamide, but not by palmitoylethanolamide, arachidonate, prostanoids, or leukotrienes (Fig. 2A). Replacement of extracellular (3K)Anandamide accumulation noVmin per milligram of protein) 15 Time (min) 0,10 0 025 0 05 0.075 0.1 0.125 1/Anandamide (nM)

Fig. 1, IA) Time course of Phlanandamide accur mulation in rat conical neurons (circles) or astrocytes (squares) at 37°C, and astrocytes at 0° to 4°C (diamonds). Results are expressed as mean = SEM of 6 to 12 independent determinations. (B and C) Lineweaver-Burk analyses of PH]anandamide accumulation (37°C, 4 min) in neurons (B) or astrocytes (C). Results are from one expenment representative of three performed in duplicate with each cell type. The firstandamide accumulation assay has been described

Na* with N-dimethylglucosamine or choline had no effect (as percentage of control: N-dimethylglucosamine, $124 \pm 12\%$; choline, $98 \pm 14\%$; mean \pm SEM, n = 6), suggesting that [H]anandamide accumulation is mediated by a Na*-independent mechanism, which has been observed with other lipids (8, 9). Moreover, inhibition of FAAH activity by creating the cells with (E)-6-(bromomethylene)tetrahydro-3-(1naphthalenyl)-2H-pyran-2-one (25 µM) or linoleyl trifluoromethyl ketone (15 µM) (13, 14) had no effect (Fig. 2, B and C) This indicates that anandamide hydrolysis did not provide the driving force for anandamide transport into astrocytes within the

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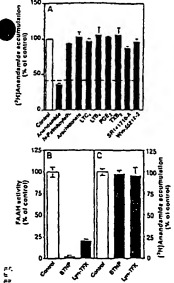


Fig. 2. (A) Selectivity of FHJanandamide accumu-ration in cortical astrocytes. Accumulation was measured after a 4-min incubation with PHJanandamide at 37°C, in the absence (control) or presence of nonradioactive anandamide (100 µM), Npalmitoylethanolamide (100 µM), arachidonate (100 µM), leukotriane C, fLTC, ; 1 µM), leukotriana B, fLTB, ; 1 µM), PGE, (100 µM), or thromboxana B2 (TXB2; 50 µM). The broken line indicales nonspecific (PH)anandamide accumulation in cells measured at 0° to 4°C (43 = 3% of total accumuit lation, which in these experiments was 43,104 = 1249 dpm per well). Results are expressed as mean ± SEM (n = 6 to 9). Effects of FAAH inho-🖆 ilors on (B) FAAH activity and (C) [PH]anandamide accumulation in cortical astrocytes. Cells were incubated for 10 min with (£)-6-(bromomethylene)tetrahydro-3-(1-naphthalenyl)-2H-pyran-2one (BTNP, 25 µM) or linoleyl influoro methylkelone (Lyn-TFK, 15 µM), and then with the same drugs plus Philanandamide for an additional 20 min. The total radioactivity in cell fipid extracts flo measure ("H)anandamide transport) (10) and radioactivity in nonesterified arachidonate to measure FAAH activity) (13) were measured separately in samples of lipid extracts prepared from the same cultures.

time frame of our experiments. Finally, the cannabinoid receptor agonist WIN-55212-2 (1 μ M) and antagonist SR-141716-A (10 μ M) also had no effect, suggesting that receptor internalization was not involved (Fig. 2A).

A primary criterion for defining earriermediated transport is pharmacological inhibition. To identify inhibitors of anandamide transport, we first examined compounds that prevent the cellular uptake of other lipids, such as fatcy acids (phloretin,

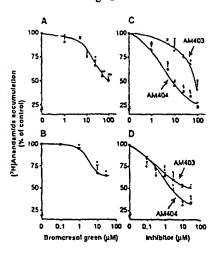


Fig. 3. Inhibition of Phljanandamide accumutation by bromcresol green in (A) astrocytes or (B) neurons: One aslensk indicates P < 0.05 and two astensks P < 0.01 Janalysis of vanance (ANOVA) followed by Bonferroni testij compared with control (H)anandamide accumulation. Inhibition of PHIanandamide accumulation by AM404 (squares) or AM403 (diamonds) in [C] astrocytes or (D) neurons. The astersk indicates P < 0.05 (paired Student's I tesi between AM404 and AM403) in all expenments, cells were incubated with the inhibitors for 10 min before the addition of PHlanandamide for an additional 4 min. Results are expressed as mean ± SEM of three to nine independent determinations.

50 μM), phospholipids (verapamil, 100 μM; quinidine, 50 μM), or POE, (bromcresol green, 0.1 to 100 µM) (15). Among the compounds tested, only bromeresol green interfered with anandamide transport, albeit with limited potency and partial efficacy (Fig. 3, A and B). Bromcresol green inhibited [3H]anandamide accumulation with an IC50 (concentration needed to produce half-maximal inhibition) of 4 µM in neurons and 12 µM in astrocytes and acted noncompetitively (16). Moreover, bromcresol green had no significant effect on the binding of [3H]WIN-55212-2 to rat cerebellar membranes (inhibition constant, K. = 22 µM), FAAH activity in rat brain microsomes ($IC_{50} > 50 \mu M$), and uptake of l'H)arachidonate or l'H)ethanolamine in astrocytes (121 ± 13% and 103 ± 12%, respectively, at 50 µM bromeresol green, n = 3) (17). The sensitivity to bromcresol green, which blocks PGE, transport, raised the question of whether anandamide accumulation occurred by means of a PGE, carrier. That this is not the case was shown by the lack of [3H]PGE, accumulation in neurons or astrocytes (18) and by the inability of PGE, to interfere with [3H]anandamide accumulation (Fig. 2A). Previous results indicating that expression of PGE2 transporter mRNA in brain tissue is not detectable further support this conclusion (9).

To search for more potent anandamide transport inhibitors, we synthesized and tested a senes of structural analogs of anandamide (19). From this screening, we selected the compound N-(4-hydroxyphenyl)arachidonylamide (AM4C4), which was both efficacious and relatively potent (Fig. 3, C and D; IC₅₀ was 1 µM in neurons and 5 µM in astrocytes). As we anticipated from its chemical structure, AM4C4 acted as a competitive

inhibitor (20), suggesting that it may serve as a transport substrate or pseudosubstrate. In contrast, at the concentrations rested AM404 had no effect on FAAH activity (IC₅₀ > 30 μ M) or on uptake of [3H]arachidonate or [3H]ethanolamine (102 \pm 4% and 96 \pm 14%, respectively, at 20 μM AM404, n = 6). Furthermore, a positional isomer of AM404, N-(3-hydroxyphenyl)arachidonylamide (AM403), was significantly less effective than AM404 in inhibiting transport (Fig. 3, C and D). These data provide pharmacological evidence for the existence of a specific anandamide transporter and suggest (i) that neurons and astrocytes may act synergistically in the brain to dispose of extracellular anandamide and (ii) that the transport systems in these two cell types may differ kinetically and pharmacologically (Fig. 1, B and C, and Fig. 3, C and D).

The identification of inhibitors allowed us to examine whether transmembrane transport participates in terminating anandamide responses mediated by cannabinoid receptor activation. Cannabinoid receptors of the CB1 subtype are expressed in neurons (21) where they are negatively coupled to adenylyl cyclase activity (22). Accordingly, in cultures of rat cortical neurons the cannabinoid receptor agonist WIN-55212-2 inhibited forskolin-stimulated adenosine 3'.5'-monophosphate (cAMP) accumulation (control: 39 ± 4 pmol per milligram of protein; 3 µM forskolin: 568 = 4 pmol per milligram of protein; forskolin plus 1 μM WIN-55212-2: 220 ± 24 pmol per milligram of protein), and this inhibition was prevented by the antagonist SR-141716-A (1 µM) (555 ± 39 pmol/mg of protein, n = 9) (23). Anandamide produced a similar effect, but with a potency (ICyo, I µM) that was 1/20 of that expected from its binding

constant for CBI cannabinoid receptors (K. = 50 nM) (1) (Fig. 4A). The transport inhibitor AM404 bound to CB1 receptors with low affinity ($K_i = 1.8 \mu M$) (19) and did not reduce cAMP concentrations when applied at 10 µM (Fig. 4B). Nevertheless, the drug enhanced the effects of anandamide, increasing the potency (by a factor of 10) and decreasing the threshold (by a factor of 1/100), an effect that was prevented by SR-141716-A (Fig. 4A). Thus, a concentration of anandamide that was below threshold when applied alone (0.3 µM) produced an almost maximal effect when applied with AM404 (Fig. 4B). Bromeresol green and AM403, which were less effective than AM404 in inhibiting anandamide transport (Fig. 3), were also less effective in enhancing the anandamide response (Fig. 4B) Furthermore, the decreases in cAMP concentrations produced by WIN-55212-2 (which stimulates CB1 receptors but is not subject to physiological clearance) or glutamate (which stimulates metabotropic receptors negatively coupled to adenylyl cyclase (24) and is cleared by a selective transporter (25)] are not affected by any of the anan-

Fig. 4. (A) Effects of AM404 on anandamide-

induced inhibition of adenytyl cyclase activity in

control neurons. The neurons were stimulated with forskolin (3 µM) in the presence of anandamide (0.001 to 3 µM; open circles), anandamide (0.001 to 3 ينكم) plus AM404 (10 ينكم) (filled

circles), anandamide (3 µM) plus SR-141716-A

(1 µM) (square), or anandamide (0.3 µM) plus

AM404 (10 µM) and SR-141716-A (1 µM) (In-

angle). (B) Effects of anandamide transport in-

hibitors on anandamide-induced inhibition of

adenylyl cyclase activity. Forskolin (FSK)-stimu-

taled neurons were incubated with AM404,

AM403, or bromcresol green (each at 10 µM) without (FSK alone) or with (FSK + anandamide)

0.3 µM anandamide. Results are expressed as

mean ± SEM of nine independent determina-

tions. One asterisk indicates P < 0.05 and two

astensks P < 0.01 (ANOVA followed by Bonfer-

roni test). (C) Effects of AM404 on the analgesic

activity of anandamide in the hot plate test. Three groups of six mice received AM404 [10]

mg/kg, intravenous), anandamide (20 mg/kg, in-

damide transport inhibitors tested (26)

These results suggest that pharmacological blockade of carrier-mediated transport protects anandamide from physiological inactivation, enhancing the potency of anandamide to nearly that expected from its affinity for CBI cannabinoid receptors in vitro. To find out whether this potentiation occurs in vivo, we tested the effects of AM404 on the antinociceptive activity of anandamide in mice. Intravenous anandamide (20 mg per kilogram of body weight) elicited a modest but significant analgesia, as measured by the hot plate test (27) (P < 0.05, Student's a test), this analgesia disappeared 60 min after injection and was prevented by SR-141716-A (Fig. 4C) (28). Administration of AM404 (10 mg/kg. intravenously) had no antinociceptive effect within 60 min of injection but significantly enhanced and prolonged anandamide-induced analgesia (Fig. 4C) (P < 0.01, Student's (test).

Our findings indicate that a high-affinity transport system present in neurons and astrocytes has a role in anandamide inactivation by removing this lipid mediator from

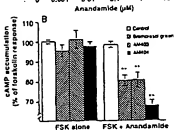
the extracellular space and delivering it to intracellular metabolizing enzymes such as FAAH (5, 6) Therefore, the identification of selective inhibitors of anandamide transport should be instrumental in understanding the physiological roles of the endogenous cannabinoid system and may lead to the development of therapeutic agents.

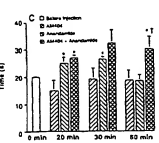
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 - Cytes (13) were prepared from rail embryos and were used after 4 to 6 days and 21 to 25 days in vitro. respectively Accumulation of PH]anandamide [22] CVImmol, New England Nuclear, Warmigton, DC was measured by moubaing the cels (six-well plates) for vanous times in Krebs buffer [136 mM NaCl, 5 mM KCl, 1,2 mM MgCl, 2 5 mM CaO₂, 10 mM glucose, and 20 mM Tatuma base; [bH 7.4], st 37°C] containing PH]anandamide in A 5 all heads. espectively Accumulation of PHJanandamide (22) 37°C) containing Philanandamide (0 45 nM, brough to 100 nM with norvadioactive anandamide), incubations were stopped by asprating the media, and cells were insed with Krebs buller containing bowne serum albumin (BSA, 0.1% w/v) and subjected to extraction with methanol and chlorolo tivity in the extracts was measured directly or after this in the case of the case o 500 nM anandamide containing 0.05 to 2.5 nM PHJanandamide We subtracted nonspecific accumulation (measured at 0° to 4°C) before determinis lonetic constants by Uneweaver-Burk analysis M Betramo and D Promelli, unpublished data.
- E. L. Barker and R. D. Blakely, in Psychopharmacology: The Fourth Generation of Progress, F. E. Bloom and D. J. Kupler, Eds. (Raven, New York, 1995), pp. 321-334, in a previous study with mixed cultures or rat conical neurons and astrocytes, a K_m of 30 µM for PHJanandamide accumulation was obtained (7). Such a high value likely resulted from the low specific radioactivity (0.3 mC/mmol) of the PHJanandamide
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 17. Displacement of PHMN-55212-2 binding (40 to 60).
- of New England Nucleary to rat coreb

80 CAMP 7 70 60 0.001 0.01 0.1 110





travenous), or anandamide plus AM404. The hol plate test (55.5°C) was performed at the times indicated, and latency to jump (in seconds) was measured before (control) and after the drugs were injected, in all groups, latency to jump before injections was 21 = 0.6 s (n = 18). A fourth group of mice received injections of vehicle alone (saline containing 20% dimethyl sulloxide), which did not affect latency to jump. One asterisk indicates P < 0.05 compared with uninjected controls (ANOVA followed by Bonlerron test).

and one cross indicates P < 0.01 compared with anandamide-treated animals (Student's I test).

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- nes (0.1 mg/ml) was determined as dememoranes (0.1 mg/mg) was determined as de-scribed JL. E. Kuster et al., J. Pramanot Epp Ther 264, 1352 (1993)) Nonspecific binding was mea-sured in the presence of 1 july novadoactive WNI-55212-2. PAM activity was measured in rat brain particulate fractions as described (1.3). The uptake of Prijarachdonate (Amosham, 200 C/mmol, 5 nm brought to 100 nM) and Prijathanolamine (Amer-sham, 50 C/mmol; 20 nM brought to 100 nM) was determined on cortical astrocytes for 4 min as of scribed (10). The control uptake for PHjarachidonale was 16729 ± 817 dpm per well and for PHjethano. ternine it was 644 ± 100 dpm per well (n = 6)
- 18. Neurons or astrocytes were incubated for 4 min at 37°C in Krebs buffer containing (PAPOE, (I).63°n M. brought to 100 nM with renradoactive PGE, 171 Cummol, New England Nuclear). After insing with Krebs buffer containing 85%, we subjected the cets to lipid extraction and counted radioactivity in the to the version and schooling in account of the version of the vers was 355 ± 28 dpm per well (n = 5).

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- The amounts of cAMP in the presence of a concentration of WIN-55212-2 below threshold (1 nM, de-26 terrined in preimmary experiments) were 96.7 = 2.5% of lorstoin stone and were not significantly affected by 10 µM AM404 (89.8 = 2.6%), 10 µM AM404 (89.8 = 2.6%), 10 µM AM405 (99.4 = 2.3%), or 10 µM bronnessed green (92.9 = 2.3%) in = 3) in the presence of a concentration of changing large. traion of gutarnate below threshold (3 µM) (24).

 CAMP concentrations were 91.5 = 2% of lorskond alone and were not significantly affected by AM404 (84 4 = 4.9%), AM400 (89.5 = 2.4%), or bromcresol

- green (84.4. = 3%) in = 3; 27. E. Fride and R. Mechautam, Eur. J. Pharmacol. 231, 313 (1993); P. B. Snidh et al., J. Pharmacol. Eup. Ther 270, 219 (1994)
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 28. The holiplate lest (55 5°C) was carried out on male Swiss med (25 to 30 g. Nossan, light following standard procedures: (F. Porreca, H. L. Mosberg, R. Hurst, V. J. Hhuby, T. F. Burks, J. Pharmacol, Erp. Ther. 230, 341 (1994)]. Arandaminie and AMADI were dissolved in 0.9% NaCl solution containing 20% dimethyl sufficiale and ripictled intavarously at 20 mouto and 10 mother, respectively. To determine 20 mg/lig and 10 mg/lig, respectively. To determine whether cannabinoid receptors participate in the ellect of anandamide, we administered anandamide. (20 mg/kg intravenously) or anadamide plus SR141716-A (2 mg/kg, subcutaneously) to living proups of sa mice each in mice that received anamamoe above. Steroy to jump increased from 21.7 ± 1.5 s to 30.7 ± 0.8 s (P < 0.05, ANOVA) 20 min after injection, in contrast, in mice that received anandamide plus SR141716-A, the latency to jump was not affected [19.6 \pm 3.1 s].
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